Course "Softwareprozesse"

**Process Improvement: CMMI**

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**Part 1:**
- Process improvement, TQM, CMMI
- The 5 CMMI Levels
- CMMI elements:
  - goals, practices
- The 22 CMMI process areas

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**Part 2:**
- Generic goals and practices
- Benefits from CMMI introduction
- Practical advice
Learning objectives

• Understand the purpose and structure of the CMMI

• Know the maturity levels and their process areas

• Learn to use CMMI as a reference framework for assessing and understanding software processes
Basic assumption of process improvement

Assumption:

When developing or building a complex product, the quality of the final product is determined to a large degree by the quality of the process used.

- This assumption is plausible for software development, because its activities tend to become so complex that even the most capable engineers cannot handle it well if the process is not well organized.
Historical notes

• ~1920: Walter Shewhart introduces Statistical Process Control at Western Electric
• ~1950-1960: William Edwards Deming pioneers continuous process improvement for industrial production in Japan
  • Similar work was performed by Joseph Juran and Armand Feigenbaum
  • This eventually led to what is now known as TQM (Total Quality Management)
  • ~1980: Deming presents his 14 points (key principles for management) and 7 deadly diseases
• 1986: Watts Humphrey and the Software Engineering Institute publish the Capability Maturity Model for Software (CMM-SW)
  • other CMMs follow (e.g. people, systems engineering) and are now combined in CMMI
Deming/TQM versus CMMI

- Deming's ideas and TQM are aimed at industrial production (manufacturing)
  - They are generic for all kinds of products
  - They pay much attention to general human factors
  - They focus on process, not product

- In contrast, CMMI is aimed at intellectual work
  - CMMI-DEV is specific to software and systems development
    - but for all kinds of software or HW/SW systems
    - There is also a CMMI-SVC for services
    - There is also a CMMI-ACQ for acquisition

- Like TQM, CMMI also pays attention to human factors
- Like TQM, CMMI also focuses on process, not product
Summary of Deming's 14 points

• Cease dependence on mass inspection to achieve quality.
  • Instead, improve the process and build quality into the product in the first place. (Feasible for production, hard for development)

• Adopt a new philosophy of cooperation (win-win).
  • Drive out fear and build trust.
  • Break down barriers between departments.

• Adopt and institute leadership for the management of people,
  • recognizing their different abilities, capabilities, and aspiration.
  • Institute training for skills.
  • Institute a vigorous program of education and self-improvement.

• Create constancy of purpose
  • for the improvement of product and service.
  • Improve constantly.

• Put everybody in the company to work to accomplish the transformation.
Basic idea of CMMI-DEV

• All high-quality software processes need to solve the same fundamental kinds of process problems

• These problems can be described in terms of
  • **process areas** and
  • **goals**

• Approaches that help solve the problems can be described in terms of
  • **practices** (describing WHAT to do, not HOW)

• These goals need to be achieved one by one.
  • Some orderings of goal-achievement are easier than others.
  • This is described by introducing **maturity levels**

• CMMI captures and represents a body of experience about useful areas, goals, practices, and goal orders.
The CMMI maturity levels

1. Initial
   - Process unpredictable, poorly controlled, and reactive

2. Managed
   - Process characterized for projects and is often reactive

3. Defined
   - Process characterized for the organization and is proactive

4. Quantitatively Managed
   - Process measured and controlled

5. Optimizing
   - Focus on continuous process improvement
Example: A process area and its goals

- One of the process areas is *Requirements Management (REQM)*
  - others are for instance *Project Planning (PP), Validation (VAL),* or *Quantitative Project Management (QPM)*
- Requirements Management is assigned to Level 2 (Managed)
  - while for instance *Quantitative Project Management* is on Level 4

Requirements Management has four goals:

- **Specific goal** SG1: Manage requirements
  - (Specific goals are particular to one process area)
- **Generic goal** GG1: Achieve Specific Goals
  - (Generic goals apply to many process areas)
- Generic goal GG2: Institutionalize a Managed Process
- Generic goal GG3: Institutionalize a Defined Process
Example (2):
A goal and its practices

**Specific goal** SG1: Manage requirements

- "Requirements are managed and inconsistencies with project plans and work products are identified."

**Specific practices** for this goal:

- SP 1.1 Obtain an Understanding of Requirements
- SP 1.2 Obtain Commitment to Requirements
- SP 1.3 Manage Requirements Changes
- SP 1.4 Maintain Bidirectional Traceability of Requirements
- SP 1.5 Identify Inconsistencies between Project Work and Requirements
Example (3): Refinements of a practice

Process area Requirements Management

- Specific goal SG 1: Manage Requirements
  - Specific practice SP 1.3 Manage Requirements Changes: "Manage changes to the req's as they evolve during the project."
    - Typical work products:
      - Requirements status
      - Requirements database
      - Requirements change requests and change impact reports
    - Subpractices:
      - Document all requirements and requirements changes that are given to or generated by the project.
      - Maintain requirements change history with change rationale and make them available to the project.
      - Evaluate the impact of requirement changes from the standpoint of relevant stakeholders.
Structure of the staged CMMI

The *staged* version of CMMI. There is also a *continuous* version.
CMMI-DEV, V1.3

Table of Contents (480 pages total)

We have just discussed ~30% of the circled region only (and will proceed with less detail).

http://www.sei.cmu.edu/library/abstracts/reports/10tr033.cfm
Required, expected, and informative elements

• An organization can have its maturity level certified
  • by a process called process appraisal or process assessment
• But only the goals described for a level are mandatory
  • the goals are *required* model elements
• while the practices are only *expected*
  • for a given organization, an alternative practice or even a non-implemented practice may be acceptable.

• Furthermore, CMMI also contains *informative* elements.
  • For instance typical work products, sub-practices, notes.
  • These may be helpful knowledge, but are purely optional.
The 22 CMMI process areas

- **Level 2: Managed**
  - Requirements Mgmt REQM
  - Project Planning PP
  - Project Monitoring&Control PMC
  - Supplier Agreement Mgmt SAM
  - Measurement and Analysis MA
  - Process and Product Quality Assurance PPQA
  - Configuration Management CM

- **Level 3: Defined**
  - Req's. Development REQD
  - Technical Solution TS
  - Product Integration PI
  - Verification VER
  - Validation VAL
  - Organizational Process Focus OPF
  - Organ'l Process Definition OPD

- **Level 4: Quantitatively Manag'd**
  - Organizational Training OT
  - Integrated Project Mgmt. IPM
  - Risk Management RSKM
  - Decision Analysis and Resolution DAR

- **Level 5: Optimizing**
  - Organizational Process Performance OPP
  - Quantitative Project Mgmt QPM
  - Organizational Performance Management OPM
  - Causal Analysis and Resolution CAR

7 + 11 + 2 + 2 process areas
Requirements Management, REQM

- **SG1** Manage requirements
  - SP 1.1 Obtain an Understanding of Requirements
  - SP 1.2 Obtain Commitment to Requirements
  - SP 1.3 Manage Requirements Changes
  - SP 1.4 Maintain Bidirectional Traceability of Requirements
  - SP 1.5 Identify Inconsistencies between Project Work and Requirements

- Often the most important/useful process area overall
Project Planning, PP

- **SG 1 Establish Estimates**
  - SP 1.1 Establish scope via a work breakdown structure
  - SP 1.2 Establish Estimates of Work Product and Task Attributes
  - SP 1.3 Define Project Life Cycle
  - SP 1.4 Estimate Effort and Cost

- **SG 2 Develop a Project Plan**
  - SP 2.1 Establish the Budget and Schedule
  - SP 2.2 Identify Project Risks
  - SP 2.3 Plan for Project-Data Management
  - SP 2.4 Plan for Project Resources

- **SG 3 Obtain Commitment to the Plan**
  - SP 2.5 Plan for Needed Knowledge and Skills
  - SP 2.6 Plan Stakeholder Involvement
  - SP 2.7 Establish and maintain the overall Project Plan
  - SP 3.1 Review Plans that Affect the Project
  - SP 3.2 Reconcile Work and Resource Levels
  - SP 3.3 Obtain Plan Commitment
Project Monitoring & Control, PMC

- **SG 1  Monitor Project Against Plan**
  - SP 1.1  Monitor Project Planning Parameters
  - SP 1.2  Monitor Commitments
  - SP 1.3  Monitor Project Risks
  - SP 1.4  Monitor Management of Process Data
  - SP 1.5  Monitor Stakeholder Involvement
  - SP 1.6  Conduct Reviews of Progress and Issues
  - SP 1.7  Conduct Milestone Reviews

- **SG 2  Manage Corrective Action to Closure**
  - SP 2.1  Analyze Issues and Determine Corrective Action
  - SP 2.2  Take Corrective Action
  - SP 2.3  Manage Corrective Action to Closure

- Bad PMC invalidates even the best PP
Supplier Agreement Mgmt, SAM

- **SG 1  Establish and Maintain Supplier Agreements**
  - SP 1.1 Determine Acquisition Type
  - SP 1.2 Evaluate and Select Suppliers Based on Criteria
  - SP 1.3 Establish Supplier Agreements

- **SG 2  Satisfy Supplier Agreements**
  - SP 2.1 Execute the Supplier Agreement
  - SP 2.2 Accept the Acquired Product
  - SP 2.3 Transition Products into Project

- Relevant not just for subcontracting but also for selecting standard software
  - such as DBMS, middleware, critical development tools etc.
Measurement and Analysis, MA

- **SG 1  Align Measurement and Analysis with Objectives**
  - SP 1.1  Establish Measurement Objectives based on Needs
    - Very important step!
  - SP 1.2  Specify Measures
  - SP 1.3  Specify Data Collection and Storage Procedures
  - SP 1.4  Specify Analysis and Reporting Procedures

- **SG 2  Provide Measurement Results**
  - SP 2.1  Collect Measurement Data
  - SP 2.2  Analyze and Interpret Measurement Data
  - SP 2.3  Store Data and Results
  - SP 2.4  Communicate Results to Stakeholders

- MA is hardly useful on Level 2
  but is an important foundation for Level 3
Process and Product Quality Assurance, PPQA

• SG 1  Objectively Evaluate Processes and Work Products
  • SP 1.1  Objectively Evaluate Process Compliance
  • SP 1.2  Objectively Evaluate Work Product Compliance
    • Work products are checked against the process description, not the project's requirements (Do not confuse this with Validation)

• SG 2  Track and Resolve Non-Compliance Issues
  • SP 2.1  Communicate and Ensure Resolution of Noncompliance Issues
  • SP 2.2  Establish Records of the Quality Assurance Activities

• Warning: PPQA will lead to 'process police' and resistance if planned processes are inadequate
  • PPQA is useful only if and where the expected process is also a sensible and suitable process
    • Less is often more
Configuration Management, CM

- **SG 1  Establish Work Product Baselines**
  - **SP 1.1** Identify Configuration Items
  - **SP 1.2** Establish a Configuration Management System
  - **SP 1.3** Create or Release Baselines
- **SG 2  Track and Control Changes**
  - **SP 2.1** Track Change Requests
  - **SP 2.2** Control Changes to Configuration Items
- **SG 3  Establish Integrity**
  - **SP 3.1** Establish Configuration Management Records
  - **SP 3.2** Perform Configuration Audits

- What this means is very project-dependent
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  - Causal Analysis and Resolution CAR
Req's. Development, REQD

- **SG 1  Develop Customer Requirements**
  - SP 1.1 Elicit Needs
  - SP 1.2 Transform Needs into Customer Requirements

- **SG 2  Develop Product Requirements**
  - SP 2.1 Establish Product and Product-Component Requirements
  - SP 2.2 Allocate Product-Component Requirements
  - SP 2.3 Identify Interface Requirements

- **SG 3  Analyze and Validate Requirements**
  - SP 3.1 Establish Operational Concepts and Scenarios
  - SP 3.2 Establish a Definition of Required Functionality
  - SP 3.3 Analyze Requirements (needed?, sufficient?, conflict?)
  - SP 3.4 Analyze Req's to Balance Needs and Constraints
  - SP 3.5 Validate Requirements with Comprehensive Methods
Technical Solution, TS

- **SG 1  Select Product-Component Solutions**
  - SP 1.1 Develop Alternative Solutions and Selection Criteria
  - SP 1.2 Evolve Operational Concepts and Scenarios
  - SP 1.3 Select Product-Component Solutions

- **SG 2  Develop the Design**
  - SP 2.1 Design the Product or Product Component
  - SP 2.2 Establish a Technical Data Package
  - SP 2.3 Design Interfaces Using Criteria
  - SP 2.4 Perform Make/Buy/Reuse Analyses

- **SG 3  Implement the Product Design**
  - SP 3.1 Implement the Design
  - SP 3.2 Develop Product Support Documentation

- TS is roughly what is commonly called *design* (in particular architecture and make/buy decisions) and *implementation*
Product Integration, PI

- **SG 1  Prepare for Product Integration**
  - **SP 1.1** Determine Component Integration Sequence
  - **SP 1.2** Establish the Product Integration Environment
  - **SP 1.3** Establish Product Integration Procedures and Criteria
- **SG 2  Ensure Interface Compatibility**
  - **SP 2.1** Review Interface Descriptions for Completeness
  - **SP 2.2** Manage Interfaces
- **SG 3  Assemble Product Components and Deliver the Product**
  - **SP 3.1** Confirm Readiness of Product Components for Integration
  - **SP 3.2** Assemble Product Components
  - **SP 3.3** Evaluate Assembled Product Components
  - **SP 3.4** Package and Deliver the Product or Component

- PI is a major aspect of what is commonly called *testing*
Verification, VER

- **SG 1** Prepare for Verification
  - SP 1.1 Select Work Products and Verification Methods
  - SP 1.2 Establish the Verification Environment
  - SP 1.3 Establish Verification Procedures and Criteria

- **SG 2** Perform Peer Reviews
  - SP 2.1 Prepare for Peer Reviews
  - SP 2.2 Conduct Peer Reviews and Identify Issues
  - SP 2.3 Analyze Peer Review Data (Conduct and Results)

- **SG 3** Verify Selected Work Products
  - SP 3.1 Perform Verification
  - SP 3.2 Analyze Verification Results

- "The purpose of Verification is to ensure that selected work products meet their specified requirements."
Validation, VAL

- **SG 1  Prepare for Validation**
  - SP 1.1   Select Products and Validation Methods
  - SP 1.2  Establish the Validation Environment
  - SP 1.3  Establish Validation Procedures and Criteria

- **SG 2  Validate Product or Product Components**
  - SP 2.1   Perform Validation
  - SP 2.2  Analyze Validation Results

- "Demonstrate that a product or product component fulfills its intended use when placed in its intended environment (such as operation, training, maintenance, support)."
  - So VAL is against user requirements (whether explicit or implicit) while VER is against product requirements and specifications
Organizational Process Focus, OPF

- **SG 1  Identify Process-Improvement Opportunities**
  - SP 1.1 Establish Organizational Process Needs and Objectives
  - SP 1.2 Appraise the Organization’s Processes
  - SP 1.3 Identify Candidate Process Improvements

- **SG 2  Plan and Implement Process Actions**
  - SP 2.1 Establish Process Action Plans
  - SP 2.2 Implement Process Action Plans

- **SG 3  Deploy Org. Process Assets, Incorporate Experiences**
  - SP 3.1 Deploy Organizational Process Assets
  - SP 3.2 Deploy Standard Processes
  - SP 3.3 Monitor the Implementation
  - SP 3.4 Incorporate Process-Related Experiences into the Organizational Process Assets

- This establishes constructive quality assurance as a potentially ongoing activity
Organizational Process Definition, OPD

- **SG 1 Establish Organizational Process Assets**
  - SP 1.1 Establish Standard Processes
  - SP 1.2 Establish Life-Cycle Model Descriptions
  - SP 1.3 Establish Tailoring Criteria and Guidelines
  - SP 1.4 Establish the Organization’s Measurement Repository
  - SP 1.5 Establish the Organization’s Process Asset Library
  - SP 1.6 Establish Work Environment Standards
  - SP 1.7 Establish Rules and Guidelines for Teams

- Lifts many Level-2 practices from project-specific forms to organization-wide standards
  - optimizes their quality, saves resources

- Like PPQA, this can lead to 'process police' and resistance if applied improperly.
  - Again, less is often more
Organizational Training, OT

- **SG 1  Establish an Organizational Training Capability**
  - **SP 1.1** Establish the Strategic Training Needs
  - **SP 1.2** Determine Which Training Needs Are the Responsibility of the Organization (as Opposed to Project or Support Group)
  - **SP 1.3** Establish an Organizational Training Tactical Plan
  - **SP 1.4** Establish Training Capability

- **SG 2  Provide Necessary Training**
  - **SP 2.1** Deliver Training
  - **SP 2.2** Establish Training Records
  - **SP 2.3** Assess Training Effectiveness
Integrated Project Mgmt., IPM

• SG 1  Use the Project’s Defined Process
  • SP 1.1  Establish the Project’s Defined Process
  • SP 1.2  Use Organizational Process Assets for Planning Project
  • SP 1.3  Establish Work Environment
  • SP 1.4  Integrate Plans
    • Extends Project Planning PP to include defined process
  • SP 1.5  Manage the Project Using the Integrated Plans
  • SP 1.6  Establish teams
  • SP 1.7  Contribute Work Products, Measurements, and Experiences to the Organizational Process Assets

• SG 2  Coordinate and Collaborate with Relevant Stakeholders
  • SP 2.1  Manage Stakeholder Involvement
  • SP 2.2  Identify, Negotiate, and Track Critical Dependencies
    • What is important for which stakeholder when?
  • SP 2.3  Resolve Coordination Issues

• IPM is most important in the context of HW+SW engineering.
CMMI Results: Effort estimation accuracy

Results: Boeing Effort Estimation

Without Historical Data
Variances between +20% to -145%
(Mostly Level 1 & 2)
(Based on 120 projects in Boeing Information Systems)

With Historical Data
Variances between -20% to +20%
(Level 3)

Reference: John D. Vu. “Software Process Improvement Journey: From Level 1 to Level 5.”
7th SEPG Conference, San Jose, March 1997.
Risk Management, RSKM

- **SG 1  Prepare for Risk Management**
  - SP 1.1  Determine Risk Sources and Categories
  - SP 1.2  Define Risk Parameters
  - SP 1.3  Establish a Risk Management Strategy
- **SG 2  Identify and Analyze Risks**
  - SP 2.1  Identify Risks
  - SP 2.2  Evaluate, Categorize, and Prioritize Risks
- **SG 3  Mitigate Risks**
  - SP 3.1  Develop Risk Mitigation Plans
  - SP 3.2  Implement Risk Mitigation Plans
Decision Analysis and Resolution, DAR

- SG 1 Evaluate Alternatives
  - SP 1.1 Establish Guidelines when to use Decision Analysis
  - SP 1.2 Establish Evaluation Criteria
  - SP 1.3 Identify Alternative Solutions
  - SP 1.4 Select Evaluation Methods
  - SP 1.5 Evaluate Alternatives Using Criteria and Methods
  - SP 1.6 Select Solutions

- The idea behind DAR:
  - A formal evaluation process reduces the subjectivity of the decision and so
  - has a higher probability of selecting a solution that meets the multiple demands of the relevant stakeholders.
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Organizational Process Performance, OPP

- **SG 1 Establish Performance Baselines and Models**
  - **SP 1.1 Select Processes to Include in Performance Analysis**
    - "What is important for us?"
  - **SP 1.2 Establish Process Performance Measures**
    - "Which measures tell us how good we are?"
  - **SP 1.3 Establish Quality and Process-Performance Objectives**
    - "How good do we need to be?"
  - **SP 1.4 Establish Process Performance Baselines**
    - "How good are we typically in process X today?"
  - **SP 1.5 Establish Process Performance Models**
    - "How does process performance change when other observable factors change?"
    - Examples: System dynamics models, Reliability growth models, Complexity models
Quantitative Project Mgmt, QPM

• **SG 1  Quantitatively Manage the Project**
  • **SP 1.1** Establish the Project’s Quality and Process-Performance Objectives
  • **SP 1.2** Compose the Defined Process
    • Select subprocesses based on performance objectives and existing performance data relative to the project requirements
  • **SP 1.3** Select the Subprocesses to be Statistically Managed
  • **SP 1.4** Select Measures and Analytic Techniques

• **SG 2  Statistically Manage Subprocess Performance**
  • **SP 2.1** Monitor Performance of the Selected Subprocesses
  • **SP 2.2** Manage Project Performance
  • **SP 2.3** Perform Root Cause Analysis
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  - Causal Analysis and Resolution CAR
Organizational Performance Management, OPM [formerly OID]

- **SG 1  Manage Business Performance**
  - SP 1.1  Maintain Business Objectives
  - SP 1.2  Analyze Process Performance Data
  - SP 1.3  Identify Potential Areas for Improvement
- **SG 2  Select Improvements**
  - SP 2.1  Elicit Suggested Improvements
  - SP 2.2  Analyze Suggested Improvements
  - SP 2.3  Validate Improvements
  - SP 2.4  Select and Implement Improvements for Deployment
- **SG 3  Deploy Improvements**
  - SP 3.1  Plan the Deployment
  - SP 3.2  Manage the Deployment
  - SP 3.3  Evaluate Improvement Effects

- In contrast to Organizational Process Focus OPF, OPM is based on quantitative management
Causal Analysis and Resolution, CAR

- **SG 1** Determine Root Causes of Defects
  - SP 1.1 Select Defect Data for Analysis
  - SP 1.2 Analyze Causes

- **SG 2** Address Root Causes of Defects
  - SP 2.1 Implement the Action Proposals
  - SP 2.2 Evaluate the Effect of Changes
  - SP 2.3 Record Causal Analysis Data

- CAR can be applied to any process quality attribute, not just product defects
The 22 CMMI process areas

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  - Project Monitoring & Control (PMC)
  - Supplier Agreement Mgmt (SAM)
  - Measurement and Analysis (MA)
  - Process and Product Quality Assurance (PPQA)
  - Configuration Management (CM)

- **Level 3: Defined**
  - Req's. Development (REQD)
  - Technical Solution (TS)
  - Product Integration (PI)
  - Verification (VER)
  - Validation (VAL)
  - Organizational Process Focus (OPF)
  - Organizational Process Definition (OPD)

- **Level 4: Quantitatively Managed**
  - Organizational Training (OT)
  - Integrated Project Mgmt. (IPM)
  - Risk Management (RSKM)
  - Decision Analysis and Resolution (DAR)

- **Level 5: Optimizing**
  - Organizational Performance Management (OPM)
  - Causal Analysis and Resolution (CAR)
Process areas are connected

E.g. the introduction to the **REQM** process area states:

- **Refer to the Requirements Development process area (REQD)** for more information regarding transforming stakeholder needs into product requirements.
- **Refer to TS** for transforming requirements into technical solutions.
- **Refer to PP** for how project plans reflect requirements and need to be revised as requirements change.

- **Refer to CM** regarding baselines and controlling changes to configuration documentation for requirements.
- **Refer to PMC** regarding tracking and controlling the activities and work products that are based on the requirements.
- **Refer to RSKM** regarding identifying and handling risks associated with requirements.

PP and CM are Level 2 areas, the others are Level 3.

Similar cross references exist in each process area.
End of part 1
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- **Each process area has its own specific goals (SG)**
  - and specific practices (SP)
Remember?: Requirements Management has four goals:

**Specific goal SG1**: Manage requirements

- Specific goals are particular to one process area

**Generic goal GG1**: Achieve Specific Goals

- Generic goals apply to many process areas

**Generic goal GG2**: Institutionalize a Managed Process

**Generic goal GG3**: Institutionalize a Defined Process

- The generic goals and practices enable the organization to institutionalize specific best practices.
  - There are only these three generic goals, GG1, GG2, GG3
    - but many generic practices
Generic goals and common features

Generic goals (GG) are realized by generic practices (GP)
These were (in CMMI v1.1) grouped into four categories:

- **Commitment to Perform (CO)**
  - Generic practices related to creating policies and securing sponsorship.

- **Ability to Perform (AB)**
  - ... ensuring that the project and/or organization has the resources it needs.

- **Directing Implementation (DI)**
  - ... managing the performance of the process, managing the integrity of its work products, involving relevant stakeholders

- **Verifying Implementation (VE)**
  - ... review by higher-level management and objective evaluation of conformance to process descriptions, procedures, and standards.
Generic practices for GG2/CO

Generic goal **GG2**: Institutionalize a Managed Process

**Generic Practices**

Commitment to Perform (**CO**):

- **GP 2.1** Establish and maintain an organizational policy for planning and performing the process.
  - Senior management should define organizational expectations for this process
Generic practices for GG2/AB

Ability to Perform (AB):

- **GP 2.2** Plan the process
  - Plan contains: process description (activities, dependencies, result requirements, quality/performance objectives), resources needed, assignment of responsibilities, training description, monitoring/measurement/review requirements, stakeholder involvement

- **GP 2.3** Provide adequate resources for performing the process
  - Funding, facilities, skilled people

- **GP 2.4** Assign responsibility and authority
  - Confirm that the people assigned to the responsibilities and authorities understand and accept them.

- **GP 2.5** Train people
  - by self-study, formalized on-the-job training, classroom training.

2.3, 2.4, 2.5 are often neglected!
Generic practices for GG2/DI

Directing Implementation (DI):  
• GP 2.6 Control work products  
  • cf. Configuration Management CM process area  
• GP 2.7 Identify and Involve the Relevant Stakeholders  
• GP 2.8 Monitor and Control the Process  
  • measure and review performance  
  • identify deviations and problems and take&track corrective action
Generic practices for GG2/VE

Verifying Implementation (VE):

- **GP 2.9** Objectively evaluate adherence of the process and address noncompliance
  - Evaluation by people external to the process
  - cf. Process and Product Quality Assurance PPQA process area
    - but PPQA is performed by the project team whereas GP 2.9 is performed by a process improvement group
    - both GP 2.9 & PPQA exist because CMMI targets large organizations

- **GP 2.10** Review process status with higher level management and resolve issues
Generic practices for GG3/CO,DI

**Generic goal GG3**: Institutionalize a Defined Process

**Generic Practices**

Commitment to Perform (CO):
- GP 3.1 Establish and maintain the description of a defined process
  - tailored from the organization's set of standard processes to address the needs of a specific instantiation

Directing Implementation (DI):
- GP 3.2 Collect Process-Related Experiences
  - e.g. effort expended for the various activities, defects injected or removed in a particular activity, and lessons learned.

No practices for: Ability to Perform; Verifying Implementation (because these are now covered by OPD process area)
Reminder: REQM Generic practices

- All of the generic practices previously seen apply to process area Requirements Management (REQM)
  - because they all apply to each Level-2 process area

- Some apply to the institutionalization of REQM on Level 2
  - those numbered GP 2.x

- others apply to REQM only when the organization moves on towards Level 3
  - those numbered GP 3.x

(And likewise for all other process areas)
In many cases, the maturity levels are too rigid
- Organizations may have good reasons for focusing on only a few process areas, rather than all of a maturity level

Therefore, there is an alternative representation of the same process areas, specific goals, and specific practices
- called the continuous representation
Staged representation

- Staged: Each *maturity level* comprises a set of process areas (whose goals must be fully reached)
Continuous representation

- Continuous: The pursuit of each goal (SG or GG) and practice has its own current \textit{capability level}
Continuous representation (2)

- Contains the same process areas and practices as staged representation
- But offers the flexibility to pursue goals in different order or intensity than prescribed by the maturity levels
  - e.g. start quantitative mgmt in one area long before Level 3 is fully addressed
- Correspondingly, the handling of generic practices is different.
- We do not discuss this any further.
Introducing CMMI

- Since the advent of CMM-SW, thousands of companies have attempted CMM-based software process improvement (SPI)

- Typical findings:
  - SPI is an expensive undertaking
  - It takes a long time (several years)
  - If it is successful, it results in improvements in many respects, e.g.
    - schedule adherence,
    - productivity,
    - SW quality,
    - customer satisfaction,
    - staff stress levels, etc.
  - It is difficult to do *fully* for smaller organisations
CMMI implementation status

- Number of organizations that report CMMI and are on level X
  - as of 2013-09

<table>
<thead>
<tr>
<th>Country</th>
<th>Appraisals</th>
<th>Maturity Level</th>
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<tbody>
<tr>
<td></td>
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<td>1</td>
</tr>
<tr>
<td>Germany</td>
<td>87</td>
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<td>21</td>
</tr>
<tr>
<td>China</td>
<td>2703</td>
<td>2</td>
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</table>
Effects from CMMI introduction

Summary of reports from conference presentations etc.

Size of improvements:

<table>
<thead>
<tr>
<th>Performance Category</th>
<th>Median</th>
<th>Number of Data Points</th>
<th>Low</th>
<th>High</th>
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<tbody>
<tr>
<td>Cost</td>
<td>20%</td>
<td>21</td>
<td>3%</td>
<td>87%</td>
</tr>
<tr>
<td>Schedule</td>
<td>37%</td>
<td>19</td>
<td>2%</td>
<td>90%</td>
</tr>
<tr>
<td>Productivity</td>
<td>62%</td>
<td>17</td>
<td>9%</td>
<td>255%</td>
</tr>
<tr>
<td>Quality</td>
<td>50%</td>
<td>20</td>
<td>7%</td>
<td>132%</td>
</tr>
<tr>
<td>Customer Satisfaction</td>
<td>14%</td>
<td>6</td>
<td>-4%</td>
<td>55%</td>
</tr>
<tr>
<td>Return on Investment</td>
<td>4.7 : 1</td>
<td>16</td>
<td>2 : 1</td>
<td>27.7 : 1</td>
</tr>
</tbody>
</table>
CMMI effects: Productivity and quality

http://www.sei.cmu.edu/cmmi/presentations/euro-sepg-tutorial/, slide 25
CMMI effects: Productivity during Level 5

- CWI: Continuous workforce innovation (P-CMM Level 5)
CMMI effects:
Source-code productivity improvement

Lockheed Martin, et al. (Kim Caputo, Beth Gramoy, Joan Weszka, Rose Whitney) "Special Intelligence from the Women in Black". SEPG. Seattle, WA, March 10, 2005.
CMMI effects: Quality improvements

- Siemens Information Systems Ltd.
  - internal document
  - 71% reduction in defect density
CMMI effects: Quality improvements

IBM Australia. "Practical Process Improvement: the Journey and Benefits."
CMMI effects: Improvements in reviews

Some Trends Experienced Along the Way

Peer Review Efficiency

Increase in peer review efficiency despite the fact that there are fewer defects to find.
CMMI effects: Quality improvements

CMMI Level 2 reduced Concorde’s post release defects by more than 40%.

On achieving Level 3 a further reduction of more than 50% was achieved.

CMMI effects: Adherence to schedule

The E-Trading team at J.P. Morgan Chase was delivering products with an average slippage of 6-8 weeks. When they achieved CMMI Level 2, the average slippage dropped to one week.

CMMI implementation steps

1 Obtain management sponsorship
   • Top management support, budget
   • Is a *Conditio sine qua non* (crucial requirement)

2 Understand the CMMI
   • Its suggestions, the ideas behind them
   • Select appropriate model(s) and representation

3 Obtain support of your organization
   • Develop/communicate business goals, rationale, costs, benefits, opportunities

4 Treat process improvement as a project
   • Form an engineering process group
   • Understand the status quo (current processes, deficiencies)
   • Sketch the target situation
   • Track progress, communicate openly
CMMI in small organizations

Why?
- Growing organizations are usually process-challenged
- Partner organizations may require some CMMI compliance

Comparison to larger organizations:
- Appraisal: relatively more expensive for a small organization
  - use simpler replacements (but not just self-assessment)
- Process definition & support: relatively even more expensive
  - prefer the continuous representation; focus on fewer areas
  - be pragmatic and inventive; learn slowly but steadily
- Deployment: may be simpler for smaller organizations!
  - people tend to be more flexible and open

http://www.sei.cmu.edu/cmmi/acss/
The Cargo Cult
The Cargo Cult
The Cargo Cult
Avoid the cargo cult!

- Many people consider CMM-based process improvement a bureaucratic monster.
- That is because many organizations tend to mistake some means for the end.
  - They think that documentation, rules, and supervising rules are process improvement.
  - Somewhat like some Melanesian island inhabitants believe that mimicking airports can bring back the prosperity they enjoyed during World War II.
DOs and DON'Ts

Senior Management:
- Don’t Treat the Level as the Goal
  - business objectives first
- **Do Pick One:** Better, Faster, Cheaper
  - be realistic, even modest
- Do Take Your Time
- Do Align the Reward System
  - reward for improvement, not only for the bottom line
- Do Lead by Example
  - Do you define your processes? And follow and improve them?

CMMI:
- Don’t Treat CMMI as the Bible
  - lots of room for improvement
  - other good sources exist
- Don’t CMMI-train the Masses
  - this just produces opposition and/or confusion
- **Become a Stronger Level 1**
  - incremental improvements are useful
  - focus on strongest pain first
- Do Use Both the Continuous and the Staged Representation
  - continuous-only may overlook important areas
  - staged-only may be too painful and slow

http://www.sei.cmu.edu/cmmi/presentations/sepg04.presentations/dos-donts.pdf
DOs and DON'Ts (2)

Measurement:
• Do Employ Basic Measures **Now!**
  • to obtain a baseline
  • schedule, effort, defects
• Don’t Collect Data You Don’t Use
  • don't create write-only DBs
  • **don't make surveys you won't act upon**
  • think in terms of return-on-investment (ROI)
• Do Enhance Data Integrity
  • invalid or undefined data is worse than no data
  • automate data collection

Process:
• Don’t Over-Engineer Processes
  • Remember Parkinson's Law
  • Rather involve many people a little (feedback/improvements)
  • Process definitions are not training materials
    • They should be rather terse
• Do Standardize Process, Not Procedure
  • What, not how
• **Do Target "Good Practice", Not "Best Practice"**
  • avoid religious wars
  • weigh consistency against cost
DOs and DON'Ts (3)

Behavioral change:

- **Do Eliminate Low-Value-Added Tasks**
  - This will also make you friends

- **Do Pilot Early and Often**
  - Theoretical considerations are often incomplete or wrong

- **Do Become a Learning Organization**
  - Reflect frequently on risks, things that work/don't work

- **Don’t Ignore the Adoption Curve**
  - target *early adopters* first
  - capture the *early majority* then
  - apply pressure to the *late majority* only then
  - eventually punish *laggards*
DOs and DON'Ts: Avoid documentation glut

How to avoid writing too much process documentation:

- Process definition **must be driven by a need**
  - Do not write more than necessary to satisfy the need
  - Write only as much as users will want to read
- Finish version 1 of each document on the day you start it
- **Focus on quality**, not quantity
  - Start with a very concise description
  - Refine the contents also by removal, not just by addition
- If a document frequently needs change, **remove detail**
- **Avoid redundancy** across documents
CMMI: So what?

- When talking about process, CMMI provides a nice framework
- We will use it as such in this course
- For each topic/proposal, we will ask:
  - What CMMI process areas are addressed?
  - How comprehensive is this proposal?
  - Can it be combined with others?
Thank you!